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7 BRUCE HORTI, et al.,  
8 Plaintiffs,  
9 v.  
10 NESTLÉ HEALTHCARE NUTRITION,  
11 INC.,  
12 Defendant.  
13

Case No. 21-cv-09812-PJH

**ORDER GRANTING DEFENDANT'S  
MOTION TO DISMISS**

Re: Dkt. No. 15

14 Defendant's motion to dismiss plaintiffs' second amended complaint ("SAC") came  
15 on for hearing before this court on June 9, 2022. Plaintiffs appeared through their  
16 counsel, Nick Suciu, III, and J. Hunter Bryson. Defendant appeared through its counsel,  
17 Timothy W. Loose. Having read the papers filed by the parties and carefully considered  
18 their arguments and the relevant legal authority, and good cause appearing, the court  
19 hereby GRANTS defendant's motion, for the following reasons.

20 **BACKGROUND**

21 This is a putative consumer class action regarding advertising of nutritional drinks.  
22 Plaintiff Bruce Horti is a resident of Concord, California. SAC ¶ 8. Plaintiff Sandra  
23 George is a resident of Adelanto, California. SAC ¶ 9. Plaintiff Jeanette Craig is a  
24 resident of Kingston, New York. SAC ¶ 10. Defendant Nestlé HealthCare Nutrition, Inc.  
25 ("Nestlé") is a Delaware Corporation with a headquarters in Bridgewater, New Jersey.  
26 SAC ¶ 11.

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1       **A. The Products**

2           Defendant makes several health drinks, including Boost Glucose Control, Boost  
3           Glucose Control High Protein, and Boost Glucose Control Max ("Boost Max"). SAC ¶ 1.  
4           Plaintiffs allege that the representations on the labels of each of these products mislead  
5           and "trick" reasonable consumers into believing that the products can prevent and treat  
6           diabetes. SAC ¶ 4. In particular, plaintiffs allege the following representations are  
7           misleading: (a) "Designed for people with diabetes"; (b) the name of the Products:  
8           "BOOST Glucose Control"; and (c) "Helps manage blood sugar." SAC ¶ 33. Boost Max  
9           does not include the representation (a) "Designed for people with diabetes." Id.

10           Plaintiffs allege they bought Boost Glucose Control drinks in retail stores. SAC  
11           ¶¶ 60-62. Each plaintiff paid an unidentified "premium price" for the drink that was "more  
12           expensive than other [unidentified] choices." SAC ¶¶ 60-62. And each plaintiff chose to  
13           purchase the drinks "based upon the Products' diabetes-related representations." SAC  
14           ¶¶ 60-62. Plaintiffs do not allege that they consumed the products, they do not describe  
15           if anything happened to them after they consumed the products, and they do not allege  
16           that they are diabetic.

17           Much of plaintiffs' complaint is dedicated to a general discussion of diabetes and  
18           other background information. SAC ¶¶ 15-59. As part of this discussion, plaintiffs  
19           concede that there is no known cure for diabetes, and that it is a condition that is  
20           managed both through "healthy eating" and taking "insulin or other medicines." SAC ¶ 21  
21           (citing Request for Judicial Notice ("RJN") Ex. B).

22       **B. Procedural Posture**

23           Plaintiffs initiated this lawsuit by complaint filed December 20, 2021. Dkt. 1. They  
24           filed the first amended complaint the same day. Dkt. 2. Pursuant to stipulation, plaintiffs  
25           filed the now-operative second amended complaint with the corrected entity name for  
26           defendant on February 4, 2022. Dkt. 9 & 11.

27           Plaintiffs assert the following claims against Nestlé: Count I: violations of  
28           California's Unfair Competition Law, Cal. Bus. & Prof. Code § 17200 ("UCL"); Count II:

1 violations of California's False Advertising Law, Cal. Bus. & Prof. Code § 17500 ("FAL");  
2 Count III: California's Consumers Legal Remedies Act, Cal. Civ. Code § 1750 et seq.  
3 ("CLRA"); Counts IV and V: New York General Business Law §§ 349 and 350 (together,  
4 "GBL"); Count VI: breach of express warranty; and Count VII: unjust enrichment. SAC  
5 ¶¶ 74-165. Plaintiffs seek to represent separate California and New York subclasses of  
6 "All persons in the [respective states] who purchased the [Boost drinks] for personal use  
7 and not for resale." SAC ¶ 64.

8 Nestlé now asks the court to dismiss the SAC in its entirety for failure to state a  
9 claim and for lack of standing. Dkt. 15. In support of the motion to dismiss, defendant  
10 requests that the court take judicial notice of certain materials. Dkt. 15-1.

#### 11 **REQUEST FOR JUDICIAL NOTICE**

12 Federal Rule of Evidence 201 permits a court to notice a fact if it is "not subject to  
13 reasonable dispute." Fed. R. Evid. 201(b). A fact is "not subject to reasonable dispute" if  
14 it is "generally known," or "can be accurately and readily determined from sources whose  
15 accuracy cannot reasonably be questioned." Fed. R. Evid. 201(b)(1)-(2). Under the  
16 incorporation by reference doctrine, the court has discretion to consider on a motion to  
17 dismiss "documents whose contents are alleged in a complaint and whose authenticity no  
18 party questions, but which are not physically attached to the [plaintiff's] pleading." Davis  
19 v. HSBC Bank Nevada, N.A., 691 F.3d 1152, 1160 (9th Cir. 2012); see also United  
20 States v. Ritchie, 342 F.3d 903, 908 (9th Cir. 2003) ("Even if a document is not attached  
21 to a complaint, it may be incorporated by reference into a complaint if the plaintiff refers  
22 extensively to the document or the document forms the basis of the plaintiff's claim.").

23 Here, defendant requests that the court take notice of Exhibit A, a reproduction of  
24 its webpages related to refunds for customers dissatisfied with the taste of Boost  
25 products. Dkt. 15-3. Though defendant contends that plaintiffs' complaint relies heavily  
26 on the Boost website, a review of the SAC in totality reveals that it relies little on the  
27 Boost website itself and rather on images from other websites or retailers. Plaintiffs have  
28 not referenced the Boost website so much that it may be considered in its entirety, and

1 defendant's request is DENIED on this basis.

2 Further, defendant's reference to the refund page of the Boost website is DENIED  
3 as moot. Defendant contends that consumer plaintiffs lack standing where a full refund  
4 was made available to them prior to suit—the remedy of a refund moots plaintiff's injury-  
5 in-fact. Savoy v. Collectors Universe, Inc., 2020 WL 4938464, at \*4 (C.D. Cal. July 21,  
6 2020). The Savoy case is much narrower than this general proposition, where a  
7 plaintiff's claim for false advertising of a customer satisfaction guarantee was deemed  
8 moot because he never attempted to utilize the refund policy prior to filing suit. Id. at \*4.  
9 But the court need not reach this argument regarding plaintiffs' standing (and thus  
10 whether to consider this exhibit) because the SAC fails on other grounds, discussed  
11 below.

12 In contrast, defendant's Exhibit B is a copy of the same informational webpage  
13 titled, "What is Diabetes?" on the CDC website that plaintiffs cite to describe diabetes in  
14 the SAC. SAC ¶¶ 18-21 n.1-4 (Dkt. 11 at 5-6). Plaintiffs' objection to the court's  
15 consideration of this material because it merely provides background on diabetes is  
16 nonsensical. Defendant cites to the material for the same purpose as plaintiffs cite to the  
17 material in their pleading, and such background information aids in assessing the  
18 "reasonable consumer" standard, an element essential to plaintiffs' claims. The court  
19 therefore GRANTS defendant's request to take notice of Exhibit B. Dkt. 15-4.

## 20 DISCUSSION

### 21 A. Legal Standards

#### 22 1. Rule 12(b)(6) – Failure to State a Claim

23 A motion to dismiss under Federal Rule of Civil Procedure 12(b)(6) tests for the  
24 legal sufficiency of the claims alleged in the complaint. Ileto v. Glock, 349 F.3d 1191,  
25 1199-1200 (9th Cir. 2003). Under Federal Rule of Civil Procedure 8, which requires that  
26 a complaint include a "short and plain statement of the claim showing that the pleader is  
27 entitled to relief," Fed. R. Civ. P. 8(a)(2), a complaint may be dismissed under Rule  
28 12(b)(6) if the plaintiff fails to state a cognizable legal theory, or has not alleged sufficient

1 facts to support a cognizable legal theory. Somers v. Apple, Inc., 729 F.3d 953, 959 (9th  
2 Cir. 2013).

3 While the court is to accept as true all the factual allegations in the complaint,  
4 legally conclusory statements, not supported by actual factual allegations, need not be  
5 accepted. Ashcroft v. Iqbal, 556 U.S. 662, 678-79 (2009). The complaint must proffer  
6 sufficient facts to state a claim for relief that is plausible on its face. Bell Atl. Corp. v.  
7 Twombly, 550 U.S. 544, 555, 558-59 (2007) (citations and quotations omitted).

8 “A claim has facial plausibility when the plaintiff pleads factual content that allows  
9 the court to draw the reasonable inference that the defendant is liable for the misconduct  
10 alleged.” Iqbal, 556 U.S. at 678 (citation omitted). “[W]here the well-pleaded facts do not  
11 permit the court to infer more than the mere possibility of misconduct, the complaint has  
12 alleged—but it has not ‘show[n]’—that the pleader is entitled to relief.” Id. at 679. Where  
13 dismissal is warranted, it is generally without prejudice, unless it is clear the complaint  
14 cannot be saved by any amendment. Sparling v. Daou, 411 F.3d 1006, 1013 (9th Cir.  
15 2005).

16 Because plaintiffs’ claims sound in fraud, their complaint must also meet the  
17 heightened pleading standard of Federal Rule of Civil Procedure 9(b). See Kearns v.  
18 Ford Motor Co., 567 F.3d 1120, 1125 (9th Cir. 2009). Rule 9(b) requires a party alleging  
19 fraud or mistake to state with particularity the circumstances constituting fraud or mistake.  
20 To satisfy this standard, the “complaint must identify the who, what, when, where, and  
21 how of the misconduct charged, as well as what is false or misleading about the  
22 purportedly fraudulent statement, and why it is false.” Salameh v. Tarsadia Hotel, 726  
23 F.3d 1124, 1133 (9th Cir. 2013) (citation and internal quotation marks omitted).

24 Review is generally limited to the contents of the complaint, although the court can  
25 also consider a document on which the complaint relies if the document is central to the  
26 claims asserted in the complaint, and no party questions the authenticity of the  
27 document. See Sanders v. Brown, 504 F.3d 903, 910 (9th Cir. 2007). The court may  
28 consider matters that are properly the subject of judicial notice, Knievel v. ESPN, 393

1 F.3d 1068, 1076 (9th Cir. 2005); Lee v. City of Los Angeles, 250 F.3d 668, 688-89 (9th  
2 Cir. 2001), and may also consider exhibits attached to the complaint, see Hal Roach  
3 Studios, Inc. v. Richard Feiner & Co., Inc., 896 F.2d 1542, 1555 n.19 (9th Cir. 1989), and  
4 documents referenced extensively in the complaint and documents that form the basis of  
5 a the plaintiffs' claims. See No. 84 Emp'r-Teamster Jt. Council Pension Tr. Fund v. Am.  
6 W. Holding Corp., 320 F.3d 920, 925 n.2 (9th Cir. 2003).

7 If dismissal is warranted, it is generally without prejudice, unless it is clear that the  
8 complaint cannot be saved by any amendment. Sparling, 411 F.3d at 1013. "Leave to  
9 amend may also be denied for repeated failure to cure deficiencies by previous  
10 amendment." Abagninin v. AMVAC Chem. Corp., 545 F.3d 733, 742 (9th Cir. 2008).

## 11 2. Rule 12(b)(1) – Lack of Article III Standing

12 The court has an ongoing obligation to ensure that it has subject matter jurisdiction  
13 such that "[i]f the court determines at any time that it lacks subject-matter jurisdiction, the  
14 court must dismiss the action." Fed. R. Civ. P. 12(h)(3). Federal courts are limited by the  
15 Constitution and Congress to only adjudicate cases involving diversity of citizenship or a  
16 federal question, or those to which the United States is a party. Mims v. Arrow Fin.  
17 Servs., LLC, 565 U.S. 368, 376-77 (2012); see also Chen-Cheng Wang ex rel. United  
18 States v. FMC Corp., 975 F.2d 1412, 1415 (9th Cir. 1992) ("Federal courts have no  
19 power to consider claims for which they lack subject matter jurisdiction."). Rule 12(b)(1)  
20 of the Federal Rules of Civil Procedure also allows a defendant to raise the defense of  
21 lack of subject matter jurisdiction by motion. The plaintiff bears the burden of establishing  
22 subject matter jurisdiction. Kokkonen v. Guardian Life Ins., 511 U.S. 375, 377 (1994).

23 A challenge to subject matter jurisdiction may be facial or factual. Safe Air for  
24 Everyone v. Meyer, 373 F.3d 1035, 1039 (9th Cir. 2004). Where the attack is facial, the  
25 court determines whether the allegations contained in the complaint are sufficient on their  
26 face to invoke federal jurisdiction, accepting all material allegations in the complaint as  
27 true and construing them in favor of the party asserting jurisdiction. Id. at 1039; Warth v.  
28 Seldin, 422 U.S. 490, 501 (1975). Where the attack is factual, however, "the court need

1 not presume the truthfulness of the plaintiff's allegations," and may review extrinsic  
2 evidence beyond the complaint without converting a motion to dismiss into one for  
3 summary judgment. Safe Air for Everyone, 373 F.3d at 1039. Once the moving party  
4 has made a factual challenge by offering affidavits or other evidence to dispute the  
5 allegations in the complaint, the party opposing the motion must "present affidavits or any  
6 other evidence necessary to satisfy its burden of establishing that the court, in fact,  
7 possesses subject matter jurisdiction." St. Clair v. City of Chico, 880 F.2d 199, 201 (9th  
8 Cir. 1989); see also Savage v. Glendale Union High Sch. Dist. No. 205, 343 F.3d 1036,  
9 1040 n.2 (9th Cir. 2003).

10 **B. Analysis**

11 **1. Whether Plaintiffs' Claims are Preempted by Federal Law**

12 "Express preemption exists when a statute explicitly addresses preemption." Reid  
13 v. Johnson & Johnson, 780 F.3d 952, 959 (9th Cir. 2015). Conflict preemption occurs  
14 when it would be "impossible for a private party to comply with both state and federal  
15 requirements," English v. Gen. Elec. Co., 496 U.S. 72, 79 (1990), or where the state law  
16 "stands as an obstacle to the accomplishment and execution of the full purposes and  
17 objectives of Congress," Hines v. Davidowitz, 312 U.S. 52, 67 (1941).

18 Defendant argues that all of plaintiffs' product labeling claims are preempted by  
19 the Food, Drug, and Cosmetic Act ("FDCA"), as amended by the Nutrition Labeling and  
20 Education Act ("NLEA"). "The [FDCA] . . . forbids the misbranding of food, including by  
21 means of false or misleading labeling." POM Wonderful LLC v. Coca-Cola Co., 573 U.S.  
22 102, 106 (2014) (citing §§ 301, 403, 52 Stat. 1042, 1047, as amended, 21 U.S.C. §§ 331,  
23 343). The NLEA provides that "no State or political subdivision of a State may directly or  
24 indirectly establish under any authority or continue in effect as to any food in interstate  
25 commerce . . . any requirement for the labeling of food . . . that is not identical to" federal  
26 requirements contained in the relevant sections. 21 U.S.C. § 343-1. "Not identical to"  
27 "means that the State requirement directly or indirectly imposes obligations or contains  
28 provisions concerning the composition or labeling of food, or concerning a food container,

1 that: (i) Are not imposed by or contained in the applicable provision . . . or (ii) Differ from  
2 those specifically imposed by or contained in the applicable provision. . . .” 21 C.F.R. §  
3 100.1(c)(4). Therefore, state labeling obligations that are “not identical to” those imposed  
4 by federal law are expressly preempted.

5 California expressly incorporates the provisions of the FDCA (as amended by  
6 NLEA) in the state’s Sherman Law. See Cal. Health & Safety Code § 110100. State  
7 laws are not preempted if they “are equal to, or substantially identical to, requirements  
8 imposed by or under” federal law. 21 C.F.R. § 808.1(d)(2). Accordingly, the NLEA “has  
9 been repeatedly interpreted not to preempt requirements imposed by state law that  
10 effectively parallel or mirror the relevant sections of the NLEA.” Lanovaz v. Twinings N.  
11 Am., Inc., No. C-12-02646-RMW, 2013 WL 675929, at \*3 (N.D. Cal. Feb. 25, 2013)  
12 (collecting cases); Clancy v. The Bromley Tea Co., 308 F.R.D. 564, 573 (N.D. Cal. 2013)  
13 (“Courts in this district have repeatedly refused to find preemption ‘where a requirement  
14 imposed by state law effectively parallels or mirrors the relevant sections of the FDCA.’”)  
15 (citations and internal punctuation omitted).

16 Defendant contests as preempted plaintiffs’ argument that the labels make “health  
17 claims” that needed to be “preauthorized by the FDA.” SAC ¶¶ 29, 45. Plaintiffs aver  
18 that their claims are not preempted because the Boost product labels at issue amount to  
19 unauthorized health claims in violation of one of NLEA’s implementing regulations, 21  
20 C.F.R. § 101.14.

21 A “health claim” is “any claim made on the label or in labeling of a food, including a  
22 dietary supplement, that expressly or by implication . . . characterizes the relationship of  
23 any substance to a disease or health-related condition.” 21 C.F.R. § 101.14(a)(1).  
24 Within the regulation, “Substance means a specific food or component of food,  
25 regardless of whether the food is in conventional food form or a dietary supplement that  
26 includes vitamins, minerals, herbs, or other similar nutritional substances.” 21 C.F.R.  
27 § 101.14(a)(2). Courts view, for example, the phrase “HEART HEALTHY/Whole grains  
28 can help support a healthy lifestyle” as a health claim because it links whole grains, a

1 substance, with heart health, a health-related condition. Hadley v. Kellogg Sales Co.,  
2 273 F. Supp. 3d 1052, 1076 (N.D. Cal. 2017) (citation omitted); see also FDA, Guidance  
3 for Industry: A Food Labeling Guide, at 81 (Jan. 2013), available at  
4 <https://www.fda.gov/downloads/Food/GuidanceRegulation/UCM265446.pdf> (“Both  
5 elements of 1) a substance and 2) a disease are present in a health claim.”). A health  
6 claim may be express or implied. Hadley, 273 F. Supp. 3d at 1074. “An implied health  
7 claim includes ‘those statements, symbols, vignettes, or other forms of communication  
8 that suggest, within the context in which they are presented, that a relationship exists  
9 between the presence or level of a substance in the food and a disease or health-related  
10 condition.’” Id. at 1074 (quoting 21 C.F.R. § 101.14(a)(1)).

11 Here, the parties do not dispute that Boost Glucose Control drinks are considered  
12 “food” under the FDCA. SAC ¶ 29. The drinks therefore must meet labeling  
13 requirements applicable to all food. See 21 U.S.C. § 343. Plaintiffs aver that the Boost  
14 Glucose Control drink is itself a “substance” as that term is defined in 21 C.F.R.  
15 101.14(a)(2). The court disagrees with the overly nuanced interpretation of the product  
16 label proffered in plaintiffs’ opposition papers. See Dkt. 16 at 22. The court also  
17 disagrees with defendant’s proposed interpretation of the regulation (see Dkt. 17 at 16),  
18 which would avoid the first half of the definition of substance, “[1] a specific food or [2]  
19 component of food.” 21 C.F.R. 101.14(a)(2) (emphasis added). The court finds that the  
20 challenged statements clearly and simply refer to the Boost products themselves as the  
21 substance that is linked to a health condition, diabetes. The court accepts that the  
22 statement “Designed for people with diabetes” refers to the Boost drink as a food when  
23 read in context with the other statements, creating a sufficient link tying the food to  
24 diabetes. The representations, collectively (a) “Designed for people with diabetes”;  
25 (b) the name of the products: “BOOST Glucose Control”; and (c) “Helps manage blood  
26 sugar” (SAC ¶ 33), establish both elements necessary to constitute an implied health  
27 claim, including (1) a substance and (2) a disease. See Hadley, 273 F. Supp. 3d at 1074  
28 (statements must be viewed in the context in which they are presented). Therefore, the

1 court concludes that the three statements on both Boost Glucose Control and Boost  
2 Glucose Control High Protein collectively constitute a “health claim” that is not  
3 preempted.

4 This assessment does not apply to one of the three products, however. The  
5 product label for Boost Glucose Control Max does not include the statement “Designed  
6 for people with diabetes.” SAC ¶ 33. That product label instead only includes the  
7 statements “BOOST Glucose Control” and “Helps manage blood sugar,” two statements  
8 that do not refer to a specific disease or health-related condition. Plaintiffs aver that  
9 “‘BOOST Glucose Control,’ is an implicit or express health claim because it purports to  
10 control a health-related condition, namely the inability to control glucose, which describes  
11 diabetes” (SAC ¶ 34), but such an interpretation suggests an inferential leap from  
12 glucose control to diabetes that falls short of specifying a health-related condition. So too  
13 with plaintiffs’ suggested inference from the statement, “Helps manage blood sugar,”  
14 where they contend that it relates to the inability to manage glucose, which describes  
15 diabetes—this leap in reasoning also goes too far and falls short of describing a health-  
16 related condition. Though managing blood sugar and controlling glucose may be  
17 important activities for persons with diabetes, these two activities do not describe a  
18 disease or health-related condition. The label of Boost Glucose Control Max does not  
19 make any health claim, and it is thus preempted because it falls outside the boundaries of  
20 the NLEA. Therefore, the court DISMISSES plaintiffs’ claims related to the third product,  
21 Boost Glucose Control Max, with prejudice.<sup>1</sup>

22 To the extent plaintiffs argue that the Boost product labels make unsubstantiated  
23 “disease claims” under 21 C.F.R. § 101.93(g)(2), their argument fails. SAC ¶ 153. As  
24 defendant highlights, that regulation does not apply because it actually deals with “dietary  
25 supplements,” a different category under the FDCA. Even if that regulation did apply,

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27 <sup>1</sup> The court does not reach the question of whether plaintiffs have standing to assert  
28 claims regarding Boost Max based on its “substantial similarity” with the other products.  
However, the fact that no plaintiff purchased Boost Max (SAC ¶¶ 60-62) certainly does  
not weigh in favor of permitting the claims related to this product to advance.

1 however, the statements on the Boost labels do not constitute “disease claims” under that  
2 regulation because the statements do not represent that “the *product itself* can cure or  
3 treat a disease.” Greenberg v. Target Corp., 985 F.3d 650, 654 (9th Cir. 2021)  
4 (emphasis added). The three statements identified by plaintiffs do not declare or suggest  
5 that the Boost products treat or cure diabetes.

## 6 **2. Whether the Product Labels Would Deceive a Reasonable Consumer**

7 Plaintiffs’ first three causes of action are brought under California statutes: the  
8 Unfair Competition Law (“UCL”), the False Advertising Law (“FAL”), and the Consumer  
9 Legal Remedies Act (“CLRA”). Cal. Bus. & Prof. Code § 17200 (UCL), Cal. Bus. & Prof.  
10 Code § 17500 (FAL); Cal. Civ. Code § 1770 (CRLA). Plaintiffs’ fourth and fifth causes of  
11 action are brought under similar New York consumer protection statutes, General  
12 Business Law sections 349 and 350.

13 “Under the consumer protection laws of California [and] New York, . . . claims  
14 based on deceptive or misleading marketing must demonstrate that a ‘reasonable  
15 consumer’ is likely to be misled by the representation.” Moore v. Trader Joe’s Co., 4  
16 F.4th 874, 881 (9th Cir. 2021); accord Consumer Advocates v. Echostar Satellite Corp.,  
17 113 Cal.App.4th 1351, 1360 (2003). “Under the reasonable consumer standard,  
18 [plaintiffs] must show that members of the public are likely to be deceived.” Williams, 552  
19 F.3d at 938. “The California Supreme Court has recognized that these laws prohibit not  
20 only advertising which is false, but also advertising which[,] although true, is either  
21 actually misleading or which has a capacity, likelihood or tendency to deceive or confuse  
22 the public.” Id. (internal quotation marks omitted) (quoting Kasky v. Nike, Inc., 27 Cal.4th  
23 939, 951 (2002)). The reasonable consumer test requires more than a mere possibility  
24 that defendant’s product “might conceivably be misunderstood by some few consumers  
25 viewing it in an unreasonable manner.” Lavie v. Procter & Gamble Co., 105 Cal.App.4th  
26 496, 508 (2003). Rather, the test requires a probability “that a significant portion of the  
27 general consuming public or of targeted consumers, acting reasonably in the  
28 circumstances, could be misled.” Id.; see also Moore, 4 F.4th at 881.

1        Courts considering the New York analogs to California's deceptive advertising  
2        claims (New York G.B.L. §§ 349, 350, plaintiffs' fourth and fifth claims here) apply the  
3        same objective assessment. Garadi v. Mars Wrigley Confectionery US, LLC, No.  
4        119CV03209RJDST, 2021 WL 2843137, at \*2 (E.D.N.Y. July 6, 2021) ("New York and  
5        California have adopted an objective definition of deception under which the alleged act  
6        must be 'likely to mislead [or deceive] a reasonable consumer acting reasonably under  
7        the circumstances.'" (citation omitted)); see also Mantikas v. Kellogg Co., 910 F.3d 633,  
8        637 (2d Cir. 2018).

9        Generally, "whether a reasonable consumer would be deceived . . . [is] a question  
10      of fact not amenable to determination on a motion to dismiss." Ham v. Hain Celestial  
11      Grp., Inc., 70 F. Supp. 3d 1188, 1193 (N.D. Cal. 2014); see Reid v. Johnson & Johnson,  
12      780 F.3d 952, 958 (9th Cir. 2015). "However, in rare situations a court may determine,  
13      as a matter of law, that the alleged violations of the UCL, FAL, and CLRA are simply not  
14      plausible." Ham, 70 F. Supp. 3d at 1193.

15        Here, plaintiffs fail to show that members of the public, particularly the targeted  
16      consumer group, are likely to be deceived by defendant's product labels. As the SAC  
17      describes, diabetes is a serious, chronic disease in which a person's ability to regulate  
18      blood sugar (glucose) is impaired. SAC ¶ 16. There is no cure, but diabetes can be  
19      *managed* "with healthy eating and being active, or your doctor may prescribe insulin,  
20      other injectable medications, or oral diabetes medicines to help manage your blood sugar  
21      and avoid complications." SAC ¶ 21; see also Dkt. 15-4. Again, plaintiffs charge that  
22      three representations on the Boost product labels are misleading: (a) "Designed for  
23      people with diabetes"; (b) the name of the Products: "BOOST Glucose Control"; and (c)  
24      "Helps manage blood sugar." SAC ¶ 33. Plaintiffs infer that these statements  
25      overpromise treatment for diabetes and underdeliver, but these statements would not  
26      lead a reasonable consumer to believe that the Boost nutritional drinks would treat this  
27      chronic disease. Reasonable consumers, particularly reasonable consumers who  
28      monitor their blood sugar, understand that consuming food, including nutritional drinks

1 like Boost, impacts blood sugar levels. See Dkt. 15-4. The product labels simply do not  
2 make the representations plaintiffs advance, where they do not represent that Boost  
3 Glucose Control will on its own treat diabetes or maintain healthy glucose levels.  
4 Plaintiffs fail to establish that persons with diabetes, as the targeted consumers of the  
5 Boost Glucose Control products, would mistake the representations for promises of  
6 treatment or even replacements for the insulin so many are prescribed. See Lavie, 105  
7 Cal.App.4th at 508 (explaining that the reasonable consumer standard is assessed from  
8 the view of “a significant portion of the general consuming public or of *targeted*  
9 *consumers*, acting reasonably in the circumstances” (emphasis added)).

10 Plaintiffs, following the reasoning in Williams v. Gerber Prod. Co., argue that the  
11 product information on the back labels of the Boost packaging cannot cure the misleading  
12 statements on the front of the package. In Williams, the Ninth Circuit determined that a  
13 reasonable consumer could be deceived by images on a fruit snack label depicting a  
14 number of different fruits, “potentially suggesting (falsely) that those fruits or their juices  
15 are contained in the product.” Williams, 552 F.3d at 939. The appellate panel rejected  
16 the argument that a misrepresentation on the front of the package could be cured by a  
17 disclaimer on the back of the package, instead concluding “reasonable consumers expect  
18 that the ingredient list contains more detailed information about the product that confirms  
19 other representations on the package.” Id. at 939-40.

20 This case, however, does not involve any misrepresentation. Though a  
21 reasonable consumer is not expected to look to the back of a product label to dispel a  
22 misleading claim on the front of a product label, see Williams, 552 F.3d at 939, plaintiffs  
23 fail to show the deception of the statements here. Plaintiffs do not contest that the  
24 product labels misstate the contents, that they misrepresent the ingredients, sugar, or  
25 carbohydrates. Where “there is no deceptive act to be dispelled and no other statement  
26 or other depiction that could mislead the reasonable consumer, then it is not plausible  
27 that a significant portion of the general consuming public could be deceived.” Ebner v.  
28 Fresh, Inc., 838 F.3d 958, 966 (9th Cir. 2016) (citing Williams, 552 F.3d at 936) (internal

1 quotation mark omitted); see also Truxel v. General Mills Sales, Inc., No. C 16-04957  
2 JSW, 2019 WL 3940956, at \*1, 4 (N.D. Cal. Aug. 13, 2019) (where “the actual ingredients  
3 were fully disclosed and it was up to the Plaintiffs, as reasonable consumers, to come to  
4 their own conclusions about whether or not the sugar content was healthy for them.”). A  
5 reasonable consumer would not be deceived by the representations on the Boost product  
6 labels.

7 The context of plaintiffs’ purchase of the allegedly misleading products only  
8 reinforces the implausibility of their claims. No reasonable consumer of the targeted  
9 consumer group would expect a novel diabetes treatment to simply appear on grocery  
10 shelves out of the blue.

11 In sum, it is not plausible that a reasonable consumer would be deceived by the  
12 Boost products labels. The court DISMISSES plaintiffs’ first five claims on this basis.

### 13 **3. Whether Plaintiffs Have Alleged a Cognizable Injury**

14 Plaintiffs may establish a cognizable injury where they did not receive the full value  
15 of a purchase by alleging that she paid a “price premium” due to the defendant’s  
16 deceptive conduct. See Izquierdo v. Mondelez Int’l Inc., 2016 WL 6459832, at \*7  
17 (S.D.N.Y. Oct. 26, 2016). However, “The bare recitation of the word ‘premium’ does not  
18 adequately allege a cognizable injury.” Naimi v. Starbucks Corp., 798 F. App’x 67, 70  
19 (9th Cir. 2019). In Naimi, the Ninth Circuit held that although the plaintiffs alleged that  
20 they paid a price premium for canned espresso, they did not allege an injury in fact  
21 because they “did not allege how much they paid for the beverage, how much they would  
22 have paid for it absent the alleged deception, . . . or any other details regarding the price  
23 premium.” Id. at 70.

24 Here, plaintiffs do not describe with any particularity how they were injured. They  
25 do not state whether they have diabetes, they do not describe whether they consumed  
26 the products, and they do not articulate any injury they suffered as a result of ingesting  
27 the Boost drinks. Plaintiffs announce that they have suffered injury based on their  
28 payment of a “premium price” for a product that did not work as advertised and that they

1 would not have paid for had they known the truth, but this is insufficient to adequately  
2 allege a cognizable injury. SAC ¶¶ 60-62. As in Naimi, plaintiffs' allegations lack any  
3 detail about the prices they paid or the differences between Boost Glucose Control  
4 products and non-premium products. Plaintiffs thus fail to make out a concrete injury.  
5 Plaintiffs lack standing for any of their claims and the SAC must be dismissed.

6 **4. Whether the Pleading Satisfies the Standards of Rule 8 and Rule 9(b)**

7 To survive the defendant's motion to dismiss, plaintiffs' CLRA, FAL, and UCL  
8 claims "must satisfy the traditional plausibility standard of Rules 8(a) and 12(b)(6), as well  
9 as the heightened pleading requirements of Rule 9(b)." Davidson v. Kimberly-Clark  
10 Corp., 889 F.3d 956, 964 (9th Cir. 2018). Satisfying this standard requires that the  
11 plaintiffs state with particularity the circumstances constituting fraud, including "the who,  
12 what, when, where, and how" of the alleged misconduct charged. Vess v. Ciba-Geigy  
13 Corp. USA, 317 F.3d 1097, 1106 (9th Cir. 2003).

14 Defendant argues that plaintiffs fall short of the necessary particularity of their  
15 reliance on the allegedly misleading product labels. See Great Pac. Sec. v. Barclays  
16 Cap., Inc., 743 F. App'x 780, 782-83 (9th Cir. 2018) (plaintiffs must plead with  
17 particularity "the 'who, what, when, where, and how' of [their] reliance."). The court  
18 agrees that plaintiffs fail to plead particularity, especially related to the "how" element of  
19 pleading under Rule 9(b). Plaintiffs allege here that each of them "relied on Nestlé'[s]  
20 diabetes-related factual representations on the Products' label[s]." SAC ¶¶ 60-62. These  
21 bare contentions fall short of describing *how* plaintiffs were led to believe that the Boost  
22 products treated diabetes or otherwise fell short of the representations on the labels.  
23 Plaintiffs offer no comparisons of these Boost products against other Boost products or  
24 other glucose-control marketed food products. Plaintiffs' CLRA, FAL, and UCL claims  
25 are not particularly pleaded, and they are therefore dismissed on this basis as well.

26 **5. Whether Plaintiffs Have Pleaded a Breach of Express Warranty**

27 "To state a claim for breach of express warranty under California law, a plaintiff  
28 must allege: (1) the exact terms of the warranty; (2) reasonable reliance thereon; and (3)

1 a breach of warranty that proximately caused plaintiff's injury." Williams v. Beechnut  
2 Nutrition Corp., 185 Cal.App.3d 135, 142 (1986).

3 Here, plaintiffs fail to identify the exact terms of a warranty. As discussed in  
4 relation to the UCL, CLRA, and FAL claims, plaintiffs have failed to identify an actionable  
5 misrepresentation as a matter of law. This claim is therefore dismissed.

6 **CONCLUSION**

7 For the reasons stated above, including plaintiffs' failures to state a claim and to  
8 plead injury-in fact sufficient to establish standing, the court GRANTS defendant's motion  
9 to dismiss. Defendant's request for judicial notice is GRANTED in part and DENIED in  
10 part, as detailed above. The court DISMISSES plaintiffs' claims related to Boost Glucose  
11 Control Max with prejudice. Plaintiffs' claims related to Boost Glucose Control and Boost  
12 Glucose Control High Protein may be amended to address the deficiencies noted in this  
13 order. Any amended pleading must be filed within 28 days from the date of this order.  
14 No additional parties or claims may be added without leave of court or stipulation of  
15 defendant.

16 **IT IS SO ORDERED.**

17 Dated: July 5, 2022

18 /s/ Phyllis J. Hamilton

19 PHYLLIS J. HAMILTON  
20 United States District Judge